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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/559,881	04/26/2000	Michael R. Schrimpf	6494.US.O2	1781
23492 759	90 06/17/2003			
STEVEN F. WEINSTOCK ABBOTT LABORATORIES 100 ABBOTT PARK ROAD			EXAMINER	
			COLEMAN, BR	ENDA LIBBY
DEPT. 377/AP6A ABBOTT PARK, IL 60064-6008			ART UNIT	PAPER NUMBER
	•		1624	
			DATE MAILED: 06/17/2003	9

Please find below and/or attached an Office communication concerning this application or proceeding.

## Office Action Summary

Application No. 09/559,881

Applicant(s)

SCHRIMPF et al.

Examiner

Brenda Coleman

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	The MAILING DATE of this communication appears	on the cover sheet with the correspondence address		
	for Reply			
A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE3 MONTH(S) FROM THE MAILING DATE OF THIS COMMUNICATION.  - Extensions of time may be available under the provisions of 37 CFR 1.136 (a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the				
mailing - If the j - If NO j - Failure - Any re	date of this communication.  Deriod for reply specified above is less than thirty (30) days, a reply within the period for reply is specified above, the maximum statutory period will apply at the reply within the set or extended period for reply will, by statute, cause the ply received by the Office later than three months after the mailing date of the patent term adjustment. See 37 CFR 1.704(b).	e statutory minimum of thirty (30) days will be considered timely.  Ind will expire SIX (6) MONTHS from the mailing date of this communication.  Be application to become ABANDONED (35 U.S.C. § 133).		
Status				
1) 💢	Responsive to communication(s) filed on Mar 24, 2	003		
2a) 🗌	This action is <b>FINAL</b> . 2b) 💢 This action	on is non-final.		
3) 🗆	Since this application is in condition for allowance e closed in accordance with the practice under Ex par	xcept for formal matters, prosecution as to the merits is te Quayle, 1935 C.D. 11; 453 O.G. 213.		
Disposit	tion of Claims			
4) 🗶	Claim(s) <u>1-38</u>	is/are pending in the application.		
4	a) Of the above, claim(s) <u>3-5, 9-18, and 29-38</u>	is/are withdrawn from consideration.		
5) 🗆	Claim(s)	is/are allowed.		
	Claim(s) 1, 2, and 19-28			
7) 💢	Claim(s) <u>6-8</u>			
8) 🗆	Claims	are subject to restriction and/or election requirement.		
	tion Papers			
9) 🗆	The specification is objected to by the Examiner.			
10)	The drawing(s) filed on is/are	a) $\square$ accepted or b) $\square$ objected to by the Examiner.		
	Applicant may not request that any objection to the di	awing(s) be held in abeyance. See 37 CFR 1.85(a).		
11)	The proposed drawing correction filed on	is: a) $\square$ approved b) $\square$ disapproved by the Examiner.		
	If approved, corrected drawings are required in reply to	o this Office action.		
12)	The oath or declaration is objected to by the Examin	ner.		
Priority	under 35 U.S.C. §§ 119 and 120			
13) Acknowledgement is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f).				
a) All b) Some* c) None of:				
1. Certified copies of the priority documents have been received.				
:	2. Certified copies of the priority documents have been received in Application No			
	application from the International Burea	cuments have been received in this National Stage u (PCT Rule 17.2(a)).		
_	ee the attached detailed Office action for a list of the			
14) X Acknowledgement is made of a claim for domestic priority under 35 U.S.C. § 119(e).				
a) U The translation of the foreign language provisional application has been received.  15) Acknowledgement is made of a claim for domestic priority under 35 U.S.C. §§ 120 and/or 121.				
•		oriority under 35 U.S.C. §§ 120 and/or 121.		
Attachment(s)  1) Notice of References Cited (PTO-892)  4) Interview Summary (PTO-413) Paper No(s)				
2) Notice of Draftsperson's Patent Drawing Review (PTO-948)  5) Notice of Informal Patent Application (PTO-152)				
3) X Information Disclosure Statement(s) (PTO-1449) Paper No(s). 5 & 6				

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DETAILED ACTION

Claims 1-38 are pending in the application.

Election/Restriction

1. Applicant's election of Group III in Paper No. 8 is acknowledged. Because applicant did

not distinctly and specifically point out the supposed errors in the restriction requirement, the

election has been treated as an election without traverse (MPEP § 818.03(a)).

2. Claims 3-5, 9-18 and 29-38 are withdrawn from further consideration pursuant to 37 CFR

1.142(b) as being drawn to a nonelected invention, there being no allowable generic or linking

claim. Election was made without traverse in Paper No. 8.

3. Claims 1, 2 and 19-28 are rejected as being drawn to an improper Markush group. The

recited compounds, while possessing a common utility, differ widely in structure and are not art-

recognized equivalents and are thus, independently distinct for the reasons set forth in the

restriction above. The Markush group represented by the term Z has variably different

definitions, rendering the claims clearly improper.

Claim Rejections - 35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

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4. Claims 25-28 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention. There is no definition in the specification for the possible additional active ingredients in claims 25-28, i.e. non-steroid anti-inflammatory agent, an opioid, a tricyclic antidepressant or an anticonvulsant.

The following is a quotation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

- 5. Claims 21, 22 and 25-28 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention. The following reasons apply:
  - a) Claims 21, 22 and 25-28 are vague and indefinite in that it is not known what is meant by an "effective amount of a compound of the formula I". It is not clear what applicants mean by an "effective amount of a compound of formula I", this does not permit the claim as written to stand alone. There is no dependency to a previous claim and therefore the claim as an independent claim must include the meanings of any and all substituents and the structure of formula I.
  - b) Claims 21 and 22 are vague and indefinite in that the claim provides for the use of claimed compounds, but the claim does not set forth any steps involved in

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determining which are the diseases capable of being ameliorated by controlling neurotransmitter release. Determining whether a given disease responds or does not respond to such an inhibitor will involve undue experimentation. Suppose that a given drug, which has inhibitor properties *in vitro*, when administered to a patient with a certain disease, does not produce a favorable response. One can not conclude that specific disease does not fall within this claim. Keep in mind that:

A. It may be that the next patient will respond. No pharmaceutical has 100% efficacy. What success rate is required to conclude our drug is a treatment? Thus, how many patients need to be treated? If "successful treatment" is what is intended, what criterion is to be used? If one person in 10 responds to a given drug, does that mean that the disease is treatable? One in 100? 1,000? 10,000? Will the standard vary depending on the current therapy for the disease?

B. It may be that the wrong dosage or dosage regimen was employed.

Drugs with similar chemical structures can have markedly different

pharmacokinetics and metabolic fates. It is quite common for pharmaceuticals to

work and or be safe at one dosage, but not at another that is significantly higher or

lower. Furthermore, the dosage regimen may be vital --- should the drug be given

e.g. once a day, or four times in divided dosages? The optimum route of

administration can not be predicted in advance. Should our drug be given as a

bolus *iv* or in a time release *po* formulation. Thus, how many dosages and dosage

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regimens must be tried before one is certain that our drug is not a treatment for this specific disease?

C. It may be that our specific drug, while active *in vitro*, simply is not potent enough or produces such low concentrations in the blood that it is not an effective treatment of the specific disease. Perhaps a structurally related drug is potent enough or produces high enough blood concentrations to treat the disease in question, so that the first drug really does fall within the claim. Thus, how many different structurally related inhibitors must be tried before one concludes that a specific compound does not fall within the claim?

D. Conversely, if the disease responds to our second drug but not to the first, both of whom are inhibitors *in vitro*, can one really conclude that the disease falls within the claim? It may be that the first compound result is giving the accurate answer, and that the success of second compound arises from some other unknown property which the second drug is capable. It is common for a drug, particularly in the treatment of pain, schizophrenia, depression etc., to work by many mechanisms. The history of psychopharmacology is filled with drugs, which were claimed to be a pure receptor XYX agonist or antagonist, but upon further experimentation shown to effect a variety of biological targets. In fact, the development of a drug for a specific disease and the determination of its biological

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site of action usually precede linking that site of action with the disease. Thus, when mixed results are obtained, how many more drugs need be tested?

E. Suppose that our drug is an effective treatment of the disease of interest, but only when combined with some totally different drug. There are for example, agents in antiviral and anticancer chemotherapy which are not themselves effective, but are effective treatments when the agents are combined with something else.

Consequently, determining the true scope of the claim will involve extensive and potentially inconclusive research. Without it, one skilled in the art cannot determine the actual scope of the claim. Hence, the claim is indefinite.

## Claim Objections

6. Claims 6-8 are objected to as being dependent upon a rejected base claim, but would be allowable if rewritten in independent form including all of the limitations of the base claim and any intervening claims.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Brenda Coleman whose telephone number is (703) 305-1880. The examiner can normally be reached on Mondays from 8:30 AM to 5:00 PM, on Tuesdays from 8:00 AM to 4:30 PM, on Wednesday thru Friday from 9:00 AM to 5:30 PM.

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The fax phone number for this Group is (703) 308-4734 for "unofficial" purposes and the actual number for **OFFICIAL** business is **308-4556**.

Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the Group receptionist whose telephone number is (703) 308-1235.

Brenda Coleman

Brenda Coleman Primary Examiner AU 1624 June 16, 2003